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(54) THE USE OF MONOMERIC/POLYMERIC MATERIALS IN THE PREPARATION OF A CURABLE COMPOSITION FOR CARTILAGE REPAIR

Die Verwendung von monomerem/polymerem Material zur Herstellung einer härtbaren Zusammensetzung für die Reparatur von Knorpeln

L'utilisation de matériaux monomériques/polymeriques pour préparer une composition durcissable destinée à la réparation de cartilage

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(56) References cited: EP-A- 0 032 249

EP-A- 0 088 845

WO-A-89/03695

· Chemical Abstracts, volume 116, no 4, 27 January 1992, (Columbus, Ohio, US), Patel, M.P.et al, "Heterocyclic methacrylates for clinical applications. II. Room temperature polymerizing systems for potential clinical use". 460, THE ABSTRACT No 28061h, Biomaterials 1991, 12 4), 649-652

Remarks:

The file contains technical information submitted after the application was filed and not included in this specification

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Description

[0001] This invention relates to the use of physiologically acceptable monomeric/polymeric materials in the preparation of a curable composition for cartilage repair by in situ curing in a human or animal body.

5 IOO22 The use of physiologically acceptable polymeric materials in the preparation of biomedical applicances such as hearing also, artificial eyes and dentures is well known, as is the use of polymeric materials as bone caments in the reflect of orthopaedics. Such polymeric materials are often used in the form of a curable composition which is initially in a fluid, seen insigned, dough-lives or other muddated from, but which cure or hardens at the temperature or use to form a rail fluid, seen insigned properties dependent on the use to which it is being put. Examples of such curable compositions are to be found in international Patent Application. NO VOB9403656 and GE Patent 2 107 341 which disclose the use of curable compositions comprising a powdered methacrylate polymer mixed with an acrylic monomer. For such applications are the preparation of hearing abits, artificial eyes and dentures which require demensional accuracy, it is important to provide compositions of low linear curing shrinkage and GB Patent 2 107 341 is directed to such an aim. For use as bone cement it is important to provide usequent failure at the bone-cement interface and, to this end.
5 WOS9030565 discloses the inclusion of a cell growth shimulant such as human growth hormone so as to increase the rate of healing of, the reample as how fractive and only ea joint of processed strends.

[003] EP-A 088,845 discloses a mixture of a monometic and polymeric acrylate or methaczylate, the monomeric component comprising a saturated oxygen containing heterocyclic saler, especially tetrahyodrufuruly asylate or methaczylate or methaczylate, as having low shinkage upon quring at room temperature, thus imparting dimensional stability. This polymeric are material is proposed for "bornedical" use, e.g. in the surgle, supplied and definite fields, especially in the man-

ufacture of hearing aids, dentures, tooth fillings and crowns. Use as a bone cement is also suggested.

[0004] EP-A 032 249 describes a mixture of tetrahydrofurfuryl methacrylate with polyethyl methacrylate as a dental

[UMU4] EP-A 032 249 describes a mixture of tetrahydrofurfuryl methacrylate with polyethyl methacrylate as a denta resin for making a temporary, but tough and adherent crown-and bridge.

[0005] M. P. Patel and M. Braden, "heterocyclic methacrytetes for clinical applications", Biomaterials 12, 649-682 (September 1991), have described compositions of tresharydorfurty methacrylate and opylethy methacrylate for producing low shrinkage materials. The implication of this paper is that these are for dertal use and for the manufacture of hearing alds. The paper person tested to the soft offer and produced the produced of the paper is the paper person tested to the soft offer and produced the produced the produced of the paper person tested to the soft offer part of the paper person tested to the soft offer part of the paper person tested to the soft offer part of the paper person tested to the soft offer part of the paper person tested to the paper person

[0006] Surprisingly, it has been found that certain curable compositions comprising a methacrylate polymer mixed with an acrylate or methacrylate monomer containing a heterocyclic group (defined below), such as terraly-directury-ly-order or methacrylate, can be employed to promote cartilage repair, with or without the addition of cell growth stimulants such as human growth hormone.

[0007] Cartilage, which differs in construction from bone, has previously been considered substantially non-regalar.

be. Attempts to repair cartilage currenty include the introduction of carbon fibres being charged in the cartilage, however such about fibres are brittle and, while a fibrous sissue forms in the implanted area, the resulting growth is not that of true cartilage composed of chordrocytes expressing when forms and perhaps and producing their own mainty components, implantation of cartilage components and hydrogel compositions have also been tried with limited success. There is therefore a grant need for a streament of cartilage orders.

[0008] According to the present invention there is provided the use of a monomer/polymer mixture in the preparation of a curable composition for introduction at or adjacent to a cartilage acquiring regain in a human or animal body to promote said repair by curing in sidu, the monomer component being selected from monomeric esters of general browns.

wherein R is a hydrogen atom or a methyl group, m is 0, 1 or 2, and X is a 3 to 6 membered heterocyclic ring and the polymer component is selected from acrylate and methacrylate polymers and copolymers thereof. Preferably X is an oxygen-containing heterocycle.

50 [0009] The monomeric ester component is preferably selected from methacrylates where X is a heterocyclic group of formula

where is 1, 2, 3 or 4. Tetrahydrofurfurf methacrylate (Ri-CH₃ m=1, n=3) is particularly preferred. These monomers to another may be admissed with other moments to control hydrophilicity, for example hydroxyphil methacrylate to increase hydrophilicity or isoborny methacrylate to decrease hydrophilicity. The polymer component is preferably a methacrylate by objecting methacrylate

[0010] The composition is suitably in the torm of a mixture of finely divided solid polymer, suitably prepared by suspension polymerisation, in liquid mornors. The composition may include initially, or have added to it at the point of use, suitable activators for the curing such as tree racical catalysts, e.g. porceide/main initiator systems. Alternatively, photo includes a cardial catalysts, e.g. powered particular systems and statistics and filters and racy openfying apents may be present. These include, for example, curinors type inhibitors in the mornomer, and/or inorganic filters to increase hardness and reduce polymerisation shrinkage, in particular hydroxypatite may be used for this purpose and to improve biocompatibility. In addition, and tibidic components such as gartantom may be added to avoid infection. Other possible therapsuitic additives include and inflammatory drugs, 19 hydrocorisonse, dewarethessone and drugs for the treatment of solicearthis when promoting sizes ergeria in a diseased joint. Other examples are artifutugal agents and animicrobial agents. Other possible additives include porcessors as collagen or decrant to increase the processly of the material or materials which function as protein onescent and control as collagen or decrant to increase the processly of the material or materials which function as protein

[0011] A particularly preferred additive is a cell growth stimulant such as those described in W089/03695, and in particular, human growth hormone. Other growth factors such as TGF-8. IGF L FDGF and FGF may be used.

[0012] The ratio of polymer to monomer component can vary dependent on the reaction time required and the consistency of composition required initially. Suitably the ratio to polymer to monomic is from 1: 10: 2: 15 weight, prefersive 12:5: 15: 17:5:1. The curing should desirably occur at body temperature and curing is desirably effected over a period of 5: big of the prefersive preferably 10: 15: minutes. The use of such compositions as biomaterials in dental, aural and confamilie fields is described in 08: 2: 107: 34: 11.

[0019] It is postulated that the ability of such cured compositions to promote tissue repair results from their ability to absorb water to an extent which allows absorption of tissue fluid from the area requiring repair, while eveiling in the tissue to provide good bonding conditions. It is postulated that, in order to be well susted for use in accordance with the present invention, the bipoclymer composition when cured should have a water uptake in the region of 5 to 30% with over a particl of its months to 2 years. It is also recognised that the low harlinkage propriets of such bipocymers combined with the slight swelling occurring with water uptake give materials which bond firmly in use, for example in cartiage, and are not readily disclosed.

[0014] The monomeripolymer imbiture is used to manufacture a composition intended for introduction at, or adjacent to damaged certilage by promote cartilage repair. His base hand not advantageous to apply the composition below, prefer services and the control of the surface of the subchondral bone in order to optimize the formation of a cartilage layer, as a result of enhanced cell printiferation and differentiation.

[0015] The invention will now be further described by way of examples.

Example 1

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[0016] A curable monomer/polymer composition was prepared by mixing polyfethy methacrylate) in powdered form (obtained from Bonar Polymers Ltd., Ret. TS: 12494, Newton Aycliffe, Co. Durham, UK. and of molecular weight 250,000) (10g) with tetrahydrotufrury methacrylate monomer (obtained from Bohm Chemie, Carmesty) (5 mortianing 2.5% v/v of N.N'-dimethyl-p-foluidine as activator. The polymer component contained 5% w/w BaSO₄ incorporated during the polymerisation process to confer andioposition.

[0017] Human growth hormone (obtained from Novo Nordisk, Denmark) was incorporated into the material by mixing 12 IU with 10g powder component prior to adding the monomer.

[0018] The composition was cast in discs of diameter 2mm at a temperature of 37°C. Elution of the growth hormone was monitored by immersion of the discs in 0.1M phosphate-buffered seline at 37°C at suitable time intervals using a specific ELISA. Figure 1 shows the <u>in ytto</u> release of human growth hormone.

[0019] The surface properties of the discs as cast were examined by scanning electron microscopy. This revealed that the cured polymer had a smooth surface, as compared to the rough surfaces obtained with more conventionally employed polymethyl methacylate.

[0000] Portions of 2mf of the curable composition prepared as above, immediately after mixing, were inserted by syringe into drilled holes in the knees of three rabbits (adult Sandy Lop, weight of least 3.5%) which were then kept unrestrained for eight months. Rapid healing and wound closure were noted and the rabbits quickly registed mobility and appetite. The tissue response at the bone-polymer interface and cartilage-polymer interface were examined. At the macroscopic level it was apported that the cartilage of eight had been polymer interface and cartilage polymer interface were examined. At the macroscopic level it was apported that the cartilage of eight had been polymer interface and cartilage.

Example 2

- [0021] A curable composition as described in Example 1 was employed. The composition was in the form of a liquid monomer containing the dispersed polymer.
- 5 (0022) A single 3 mm diameter defect was drilled into the intercondylar notor of the articular cartilage of 18 mature. Sandy Lop rabbits. Into each defect was inserted 0.15 ml of the curable composition containing human growth hormone. It? International units per 5g polymer powder) into the subchondral bone below the area of removed cartilage. Plain curable composition without any growth hormone was inserted into a similar cartilage defect in the contralateral limb. Rabbits were accrificant at 3.6 g and 12 weeks and 8 months.
- 10 [0023] <u>Histology</u>. At each time interval, excess bone was removed from the femoral conclyles before fixation in 2% paraformaldehyde, and 0.5% glutardehyde, at 4°C, for to 48 hours. Each specimen was decalcified in neutral EDTA at 4°C prior to either low temperature (4°C) delignation and was embedding, or frozen section preparation.
- [0024] <u>Cryosectionins</u>. At each time interval decalcified specimens were frozen in cryomountant using liquid nitrogen. These were cryostat sectioned at .20°C, whilst the polymer was held in place with double-sided selfotape, sections were then mounted on glass slides and stained.
 - [0025] Immunolocalisation of collagen hose II. chondroitin 4 subhate and chondroitin 6 subhate. Dewaxed sections were chondroitinase digested (0.25 IUI/ml) for one hour at 38°C to reveal the epitopes before immunolocalisation. Selected primary monoclonal antibodies were used individually and a rhodamine conjugated anti-murine serum was applied to each section, after appropriate wash steps between stages. Non-immune mouse serum was applied as con-
- 20 tol to all immunolocalisations and pre-ebsorbed anti-collegen type II monoclonal was used as an extra control for the localisation of collagen type II which is specific to arriage.
 [0026] <u>Electron Microscopy</u> At each time Interval, excess bone was removed from the femoral condyles before fine
- trimming to the area of delect repair, before fixation with 25% glutarablehyde in sodium caccolytate buffer at 47°C, for a minimum 48 hours. Specimens were post fixed in 1% certain tetroxide for 2 hours before dehydration through a methanol series into provilene oxide and embedding in Souri's resident.
- [0027] The results observed were as follows:
 - [0028] <u>Macroscopic findings</u>. None of the rabbits died, nor was there any evidence of infection. The rabbits were housed in group pens which allowed freedom of movement i.e. running, jumping, standing on hind legs. The animals showed no sign of discomfort and all enjoyed full mobility.
- 30 In most rabbits the knee joints showed a white plistening cartilage-like tissue resembling the normal surrounding articular cartilage. There appeared to be a good overgrowth of cartilage over the polymer within the defect. In three tabbit knees the tissue covering was incomplete; histological observations revealed that the polymer had been set above the level of the subchordral bore in the cartilage defect. Since cartilage cannot grow through the polymer, it is therefore important that the polymer is set at the right level or low good resurration.
- 35 [0022] Two rabbits were legst for a longer study (eight morths). The joints remained functional throughout the study period. The histology revealed that the new carsinger emained intact but he density of the metrix had still not achieved that of the original cartilage. There were still a mixed population of cells and areas of librous and chondrogenic regions. The subchondrab lone had remodeled and in it he polymer became surrounded by very dense coalgen.
- [0030] Figzen sections, Cryostat sectioning allowed visualisation of the intact polymer-issue interface, hence the obtacellular matrix components of the lissue growing over the polymer three weeks after surgery were characterised. Histologically a variety of itsues were observed. Most prominant in the early stages was the observation of a highly cellular fibrous tissue. A thin layer of synovial appearance separated the new issue covering the polymer from the inter-condylar space. Broys picules appeared to be associated with areas of new issue immediately adjacent to the polymer strace. Above this interface the fibrous layer contained areas of rounded cells in a metachromatically stained metrix believed to be chrondropein roundules.
- [0031] Low temperature wax embedded tissue sections. Immunolocalisation of collagen type II within the cartiaginus nockles confirmed the chordrogenic phenotype of these areas of the tissue. Immunolocalisation studies also demonstrated an elaboration of chordroff in 4-subhate and chondroff in 5-suphate glovaminoglycan side chains both in the fibrous tissue and in the regions of chordrogenic nockles. The varieties of cell phenotypes within the layer cov-
- 50 ering the polymer were shown histologically and by immunolocalisation of their matrix molecules during the first welve weeks after implantation. [0032] Transmission electron microscopy of the trans-polymer tissue layer after 8 months of implantation showed the
 - [U032] <u>Itansmission electron microscopy</u> of the trans-polymer tissue layer after 8 morths of implantation showed the rounded appearance of cells within a proteoglycan rich matrix, indicating the chondrogenic nature of the tissue. The presence of chondron at the cartilage and bone interface was noted.
- 55 [0033] <u>Growth hormone incorporation</u>. It was shown that the cured polymer system was a good vehicle for the release of growth hormone. Morphological comparisons were made between the tissue covering the growth hormone and plain polymer.
 - [0034] Figures 2 to 6 illustrate the above findings as follows:

Figure 2

[0035] After 3 weeks of implantation a fibrous tissue layer (f) had grown over the polymer (P) surface. The polymer (P) had been inserted into a sub-chondral defect in the bone (B) below the level of the remaining cartilage (C).

[0036] Decalcified tissue embedded in wax. Section is stained with Methylene blue-Azur II.

Figure 3

[0037] After 6 weeks of implantation wo zones of repair tissue were observed. Bony spicules (b) and nodules contraining chandrocytes (arrowed) are seen in the zone immediately above the polymer (P) surface. Where the defect has been made in the carrillage (c) a issue layer similar in appearance but less organised than normal carrillage (c) has formed above the bory zone. Original bone is denoted 8, and the original carrillage is denoted 0. [0038] Deachiffed tissue embodded in wax. Section is stained with Methane blue. Azur II.

15 Figure 4

[0039] After 9 weeks of implantation there is more new bone (b) above the polymer (P) surface. There are nodules containing chondrocytes (arrow) in the borry layer. The new cardiage (c) is disorganised. Original bone is denoted B, and the original cardiage is denoted C.

20 [0040] Decalcified tissue embedded in wax. Section is stained with Methylene blue-Azur II.

Figure 5

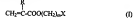
[0041] By transmission electron microscopy clusters of chondrocytes are seen to be contained within a collagenous is basket (co). These are referred to as chondrons and are structures normally observed in the deep zones of mammalian cardiage. The normal extracellular matrix is denoted M.

Figure 6

30 [0042] The cells within the chondron (Co) appear to be actively synthesising cell products indicated by the enormous amount of endoclasmic reticulum (ER). The cell nucleus is denoted N and the normal extracellular matrix is denoted M.

Claim

35 1. The use of a monomer/polymer mixture in the preparation of a curable composition for introduction at or adjacent to a cartilage site requiring repair in a human or animal body to promote said repair by curing in situ, the monomer component being selected from monomeric esters of central formula.



where R is a hydrogen atom or a methyl group, m is 0, 1 or 2 and X is a 3 to 6 membered heterocyclic ring and the polymer component is selected from acrylate and methacrylate polymers and copolymers thereof.

- 2. A use according to Claim 1, wherein X in the monomer component of formula I is an oxygen-containing heterocycle.
- 3. A use according to Claim 2, wherein the monomer component is tetrahydrofurfuryl methacrylate.
- 4. A use according to Claim 1, 2 or 3, wherein the polymer component is poly(ethyl methacrylate).
- A use according to any one of the preceding claims in the preparation of a curable composition having a polymer to monomer ratio of 1:1 to 2:1 by weight.
- A use according to any one of the preceding claims, in the preparation of a curable composition further comprising one or more components selected from antibiotic and therapeutic agents, antifungal and antimicrobial agents, porosogens, protein carriers and cell growth stimulants.

7. A use according to Claim 6, wherein the, or one of the, additives is human growth hormone.

Patentansprüche

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5 1. Varwendung einer Monomer/Polymer-Mischung zur Herstellung einer haftbaren Zusammensetzung zum Errichten bei oder in Nachberschaft zu einer Knorpstelled, die einer Repearatte badert, in einem menschlichen oder tierlichen Korper, um die Reparatur durch in situ erfolgendes Härten zu tordern, wobei die Monomerkomponente ausoweithst aus ernoomeren Esten der allebereinen Formel 1.

$$CH_2=C-COO(CH_2)_mX$$
 (I),

in der R ein Wasserstoffatom oder eine Methylgruppe ist, m 0, 1 oder 2 ist und X ein 3- bis 6-gliedriger heterocyclischer Ring ist und die Polymerkomponente ausgewählt ist aus Acrylat- und Methacrylatpolymeren und Copolymeren davon.

- Verwendung nach Anspruch 1, wobei in der Monomerkomponente der Formel IX ein Sauerstoff enthaltender Heterocyclus ist.
 - 3. Verwendung nach Anspruch 2, wobei die Monomerkomponente Tetrahydrofurfurylmethacrylat ist.
- 25 4. Verwendung nach Anspruch 1, 2 oder 3, wobei die Polymerkomponente Poly(ethylmethacrylat) ist.
 - Verwendung nach einem der vorangehenden Ansprüche zur Herstellung einer h\u00e4rtbaren Zusammensetzung mit einem Gewichtsverh\u00e4ltnis von Polymer zu Monomer von 1:1 bis 2:1.
- Verwendung nach einem der vorangehenden Ansprüche zur Herstellung einer h\u00e4rbaren Zusammensetzung, die ferner eine oder mehrere Komponenten, ausgewahlt aus antibiotischen und therapeutischen Nitteln, fungiziden und antimirckobleiten Mitteln, Prosisitätsbildnern, Proteinträgern und Zellwechstumsstimutantien, umfalls.
 - 7. Verwendung nach Anspruch 6, wobei das oder eines der Additive menschliches Wachstumshormon ist.

Revendications

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Utilisation d'un mélange monomère polymère dans la préparation d'une composition durciseable destinés être introduite à Perplacement d'un cartilage ou à son voisinese, nécessitant une réparation dans un corps humain ou animal afin de tavoriser ladite réparation par durcissement in situ, le composant monomère de tormule definérale !

dans laquelle R est un atome d'hydrogène ou un groupe méthyle, m est 0, 1 ou 2 et X est un noyau hétérocyclique a 3 - 6 éléments et le composant polymère est choisi parmi des polymères acrylates et méthacrylates et les copolymères de ceux-ci.

- Utilisation selon la revendication 1 dans laquelle X, dans le composant monomère de formule I, est un hétérocycle contenant de l'oxygène.
- Utilisation selon la revendication 2 dans laquelle le composant monomère est le méthacrylate de tétrahydrofurfurryle.

- 4. Utilisation selon la revendication 1, 2 ou 3 dans laquelle le composant polymère est le poly (méthacrylate d'éthyle).
- Utilisation selon l'une quelconque des revendications précédentes dans la préparation d'une composition durcissable ayant un rapport polymère : monomère de 1 :1 à 2 :1 en poids.

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- Utilisation solon l'une quélconque des revendications précédentes dans la préparation d'une composition duriés abble comprenant en outre un ou plusieurs composants choisis parmi les agents ambitiques et tréspectulques, les agents artiflongiques et antimicrobiens, les porosogènes, les porteurs protéiques et les stimulateurs de croissance cellulation.
- 7. Utilisation selon la revendication 6, dans laquelle les additifs, ou l'un de ceux-ci, est l'hormone de croissance

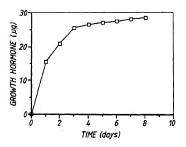


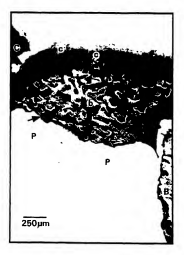
Fig.1

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F<u>ig</u>.2

9



F<u>ig</u>.3



F<u>i</u>g.4

